



**University of
Zurich**^{UZH}

**Zurich Open Repository and
Archive**

University of Zurich
University Library
Strickhofstrasse 39
CH-8057 Zurich
www.zora.uzh.ch

Year: 2018

Disturbance reverses classic biodiversity predictions in river-like landscapes

Harvey, Eric ; Gounand, Isabelle ; Fronhofer, Emanuel A. ; Altermatt, Florian

Abstract: Global analyses of biodiversity consistently reveal recurrent patterns of species distributions worldwide. However, unveiling the specific mechanisms behind those patterns remains logistically challenging, yet necessary for reliable biodiversity forecasts. Here, we combine theory and experiments to investigate the processes underlying spatial biodiversity patterns in dendritic, river-like landscapes, iconic examples of highly threatened ecosystems. We used geometric scaling properties, common to all rivers, to show that the distribution of biodiversity in these landscapes fundamentally depends on how ecological selection is modulated across space: while uniform ecological selection across the network leads to higher diversity in downstream confluences, this pattern can be inverted by disturbances when population turnover (i.e. local mortality) is higher upstream than downstream. Higher turnover in small headwater patches can slow down ecological selection, increasing local diversity in comparison to large downstream confluences. Our results show that disturbance-mediated slowing down of competitive exclusion can generate a specific transient signature in terms of biodiversity distribution when applied over a spatial gradient of disturbance, which is a common feature of many river landscapes.

DOI: <https://doi.org/10.1098/rspb.2018.2441>

Posted at the Zurich Open Repository and Archive, University of Zurich

ZORA URL: <https://doi.org/10.5167/uzh-172063>

Journal Article

Accepted Version

Originally published at:

Harvey, Eric; Gounand, Isabelle; Fronhofer, Emanuel A.; Altermatt, Florian (2018). Disturbance reverses classic biodiversity predictions in river-like landscapes. *Proceedings of the Royal Society of London, Series B: Biological Sciences*, 285(1893):1-9.

DOI: <https://doi.org/10.1098/rspb.2018.2441>

1 **Title:** Disturbance reverses classic biodiversity predictions in river-like landscapes

2 **Running head:** Disturbance in river-like landscape

3 **Authors:** Eric Harvey^{1,2,3*}, Isabelle Gounand^{1,2*}, Emanuel A. Fronhofer^{1,2,4}, and Florian

4 Altermatt^{1,2}

5 *These authors contributed equally to this work

6 **Corresponding Author:** eric.harvey@umontreal.ca

7 **Affiliations:**

8 ¹Department of Evolutionary Biology and Environmental Studies, University of Zurich,

9 Winterthurerstrasse 190, CH-8057 Zürich, Switzerland.

10 ²Eawag, Swiss Federal Institute of Aquatic Science and Technology, Department of

11 Aquatic Ecology, Überlandstrasse 133, CH-8600 Dübendorf, Switzerland

12 ³Department of Ecology and Evolutionary Biology, University of Toronto, M5S 3B2,

13 Canada.

14 ⁴ISEM, Université de Montpellier, CNRS, IRD, EDPHE, Montpellier, France.

15 **Statement of authorship:** EH, IG, EAF and FA designed the research; IG and EAF

16 designed the model; IG programmed and ran the model, analyzed the simulation data

17 with support from EAF and produced the figures; EH conducted the lab experiment with

18 support from IG, EAF and FA, processed the experimental data with support from IG,

19 and carried out the analysis of experimental data; all authors participated in results

20 interpretation; EH wrote the first draft of the manuscript; All authors significantly

21 contributed to further manuscript revisions. EH and IG contributed equally to this work.

22 **Keywords:** Meta-community, Dendritic Networks, Biodiversity, Patch size, Perturbations

23

Abstract

Global analyses of biodiversity consistently reveal recurrent patterns of species distributions worldwide. However, unveiling the specific mechanisms behind those patterns remains logistically challenging, yet necessary for reliable biodiversity forecasts. Here, we combine theory and experiments to investigate the processes underlying spatial biodiversity patterns in dendritic, river-like landscapes, iconic examples of highly threatened ecosystems. We used geometric scaling properties, common to all rivers, to show that the distribution of biodiversity in these landscapes fundamentally depends on how ecological selection is modulated across space: while uniform ecological selection across the network leads to higher diversity in downstream confluences, this pattern can be inverted by disturbances when population turnover (i.e., local mortality) is higher upstream than downstream. Higher turnover in small headwater patches can slow down ecological selection, increasing local diversity in comparison to large downstream confluences. Our results show that disturbance-mediated slowing down of competitive exclusion can generate a specific transient signature in terms of biodiversity distribution when applied over a spatial gradient of disturbance, which is a common feature of many river landscapes.

Introduction

Local species diversity in dendritic, river-like landscapes is generally expected to be higher in larger, more connected downstream confluences than in upstream headwaters [1–6]. Previous theoretical, comparative and experimental studies have all emphasized the importance of dispersal along the network structure of the landscape as a key driver of this diversity pattern, regardless of the specific local drivers of community dynamics such as ecological drift [3,7] or selection [8]. The pattern of lower local diversity in upstream habitats compared to downstream confluences (hereafter ‘classical pattern’), however, is not ubiquitous in natural river systems: recent empirical studies [9,10] have documented that diversity patterns can be completely reversed, with higher diversity in upstream headwaters rather than in downstream confluences (hereafter ‘reversed pattern’). These contrasting empirical patterns, especially the reversed one, and the transition from one to the other remain insufficiently understood and are currently not accounted for by any theoretical or experimental work. Given that river ecosystems support roughly 10% of all animal species [11], it is unfortunate that we are still lacking a general understanding of the processes driving the patterns of diversity in these ecosystems. A better understanding of the dominant mechanisms that generate contrasting diversity patterns is essential if we hope to forecast effects of global change and possibly design management strategies to counter current trends of erosion of biodiversity and ecosystem services in river ecosystems worldwide [12–14].

Riverine landscapes are structured by spatial flows of organisms and resources [1,15]. They also inherently display strong environmental gradients, both in terms of resources (habitat size / land use influence) and disturbances (current flow, erosion) [1,16]. While

previous theoretical work has primarily focused on dispersal-related mechanisms [3,17,18], the latter intrinsic characteristics might be key to explain contrasting diversity patterns in such spatially structured systems where habitat size scales with position in the landscape [4,15,16,19]: Small headwater patches (“patch” *sensu* [20]) can be found across a wide range of landscape connectivity, but fewer and larger downstream patches are inherently more connected [16]. The Theory of Island Biogeography [21,22] provides explanation for one of the most established and universal ecological patterns stating that smaller and more isolated patches suffer higher extinction rates and lower immigration rates eventually leading to lower diversity than in larger and more connected patches [21–25]. In such a scenario, competitive exclusion due to ecological selection occurs more rapidly in small headwater communities because, in finite populations, extinction thresholds are reached earlier in smaller populations than in large ones. Consequently, all else being equal, at equilibrium, smaller patches should always contain lower diversity. In that context the ‘reversed pattern’ can only occur if (i) all else is not equal and smaller headwater patches contain for instance higher environmental heterogeneity or (ii) the equilibrium state is reached more slowly in smaller patches. The former represents the obvious case where headwater patches might contain higher proportion of micro-habitats providing higher niche dimensionality and/or refugia from competition and predation. Here we focus on the latter case, where disturbances have the potential to keep smaller patches away from their equilibrium community composition. Given that upstream headwaters are less buffered from disturbances relative to larger downstream confluences, they are naturally more affected by disturbances such as desiccation or fluctuation in contaminants [26,27] and are increasingly affected by human impacts such

as mountain top mining [28]. Therefore, our study addresses an especially likely and increasingly widespread scenario.

Disturbances are often seen as an important driver of community dynamics by constraining the growth of more competitive species, which slows down competitive exclusion, and ultimately can favour transitory coexistence periods that are long enough to be empirically observed [29,30]. If disturbances impact smaller patches more strongly than larger patches, population turnover (here referring to local mortality, its underlying mechanism) will be increased, which can potentially lead to higher species richness in these smaller headwater patches than in larger downstream patches. More specifically, under the classical biodiversity pattern, competitive exclusion by ecological selection is expected to occur faster in smaller patches relative to larger patches because of lower population sizes (i.e., closer to extinction threshold) and potentially higher encounter rates among more spatially confined species (i.e., stronger interaction strengths – see [31]). In the presence of disturbances, however, ecological selection could be slowed down, more so in smaller than in larger patches, potentially reversing the spatial biodiversity pattern. Despite strong empirical evidence on the role of disturbances in driving biodiversity patterns in riverine ecosystems, via its impact on competitive dynamics [2,32–36], it remains highly challenging to investigate this hypothesis directly in the field due to multiple interacting factors and lack of replication. Therefore, to get a mechanistic understanding, we first develop a general mathematical model, and then test the main predictions in well-established ecological microcosm landscapes [37,38].

Specifically, we here investigate the processes underlying the distribution of biodiversity in river-like landscapes and focus on characteristic patch size distributions

and disturbances (i.e., local mortality). In natural riverine landscapes, patch size distribution, network configuration, and disturbances are intrinsically linked, and cannot be disentangled in a causal approach [4,16,39]. Thus, here we combined theory and experiments to untangle these naturally interlinked factors. We first show theoretical results from a spatially explicit Lotka-Volterra competition metacommunity model including dispersal along dendritic networks, demographic stochasticity, and patch size-dependent mortality, that is, higher local mortality in smaller patches (see full details in Methods). We secondly tested experimentally the model's main predictions, using dendritic protist microcosm landscapes [38] of the same network topologies (see Methods and Fig. 1). Our work shows that patch size-dependent mortality, and its effect on ecological selection can be a key mechanism shaping transient biodiversity patterns in entire river-like landscapes.

Material and Methods

Model Simulation

To investigate the mechanisms underlying biodiversity distributions in dendritic, river-like networks in relation to their intrinsic non-random patch connectivity/size pattern, we first used general simulations of a Lotka-Volterra competition model for 10 species, in which the temporal variation in abundance of species i located in patch x ; N_{ix} , is described by:

$$\frac{dN_{ix}}{dt} = r_i N_{ix} \left(1 - \frac{\sum_{j=1}^n \alpha_{ij} N_{jx}}{K_{ix}} \right) - \sum (d_x N_{ix} - d_z N_{iz}) - m_x N_{ix}$$

where r_i is the intrinsic growth rate, α_{ij} the per capita effect of species j on species i (competition coefficient), K_{ix} the carrying capacity of species i in patch x (K is higher in larger patches), d_x the dispersal rate from patch x , and m_x the patch size-dependent mortality rate (population turnover). Demographic stochasticity is added according to a standard method [40] with the following term $+a\rho\sqrt{N_{ix}}$. The parameter a modulates the magnitude of this drift and ρ is the random factor sampled from a Gaussian distribution. Parameters are set as in Giometto *et al.*, 2013 [40] to produce a white noise preventing artificial persistence of very small populations (see Table 1). Species traits were parameterized in a very simple and general way, independent of our experimental organisms, to preserve the generality of our model predictions. To limit sources of variations, r , K and d are the same for all species (see values in Table S1 in Supporting Information) and we implement species differences via competition coefficients only. Competition coefficients are drawn from a Gaussian distribution ($\mu = 2$, $\sigma = 0.25$) with all negative coefficients multiplied by -1 to obtain a purely competitive community. The dispersal term is a sum function of the differences between immigration from patch x and each emigration from patch z to which it is connected according to the specific topology of the dendritic landscape. We scale dispersal rates with patch size to make it comparable to the experimental set-up where exchanging a constant volume between two patches was the most feasible dispersal mode. Then dispersal rate was implemented as $d_x = \delta_V/V_x$, with δ_V the fixed volume dispersed by unit of time from a patch x and V_x the volume of the patch x . In our simulations we vary δ_V to vary dispersal rates. Our results are, however, robust to change in dispersal mode whether the dispersal rate is a constant rate either per edge or per patch (see Appendix A and Fig. S1). Moreover, while we

considered only bi-directional dispersal to accommodate logistical constraints on the experimental design, we also conducted a sensitivity analysis, which attests that the results are robust to a strongly biased downstream dispersal (Appendix A and Figs. S2-S3). As a minimal assumption to implement patch size dependency in turnover, we make local mortality rates follow a linear negative relationship with patch size (see sensitivity analysis below for the values). We report results with dispersal and local mortality rates set at intermediate levels ($\delta_V = 0.05$, and m are 0.1, 0.0868, 0.064 and 0.01 respectively from smaller to larger patches) in the first part of the results (Figs. 1–3), while we explore further the parameter space with multiple combinations of those two rates in Fig. 4 (see below for more details).

For the landscapes, we use river-like networks generated from five different space-filling optimal channel networks [41] known to reproduce the scaling properties observed in real river systems [4,42]. Optimal channel networks are built under the assumption that drainage network configurations always minimize total energy dissipation, and the empirical observation that river network properties are scale-invariant (i.e., fractal, see Rinaldo *et al.*, 2006 [42]). In order to also be able to use the same landscapes in the experiment, a coarse-graining procedure is used to reduce the 5 generated constructs to equivalent 6x6 patch networks, preserving the characteristics of the original three-dimensional basin (see Fig. 1, for details see appendix A in Carrara *et al.*, 2014 [4]), but having a total patch size that was experimentally feasible. These 6x6 patch networks contained 36 patches of four different volumes (V): 7.5, 13, 22.5, and 45 mL (see Fig. 1). The ratios between those different volumes are based on the known relationship between the size of a channel reach and the landscape-forming discharge at-a-station within the

catchment (see [16] and appendix A in Carrara *et al.* 2014 [4] for more details). These volume values are also used for patch sizes V_x in the model.

Using these settings, we test the effect of patch size-dependent mortality on biodiversity patterns by contrasting simulations with and without disturbance. We have two levels of replication: we replicate our simulations over the 5 different dendritic landscapes to assess the independence of the results from specific network topologies, and 10 random realisations of the community matrix (replicate communities), leading to 50 simulations in total per scenario (local mortality settings at a given dispersal rate). Simulations always start with all species already present in all patches and with the same absolute abundance set to 200. The results are robust to starting conditions with same relative abundance (scaled to patch size). Simulations of community dynamics are run with the Runge-Kutta Cash-Karp method with adaptive step-size control (GSL 1.15) to optimize the accuracy of numerical integration in continuous time. Simulations are stopped at 400 units of time, which is sufficient to observe competitive exclusion (see Fig. 3 and Fig. S4). A species for which abundance falls below 10^{-3} is considered extinct and its abundance set to 0. The results are robust to the decline or suppressing of this threshold but we keep it to optimize the computation time.

In addition, we also explore the persistence of the different observed diversity patterns as a function of the magnitude of dispersal and the strength of the disturbance (local mortality) vs. patch size relationship (see Figs. 4, S5 and S6). We explore 6 levels of dispersal with $\delta_V \in \{0, 0.05, 0.1, 0.25, 0.5, 1\}$ and 21 mortality rate combinations. We used all mortality rate combinations with $m \in \{0, 0.01, 0.05, 0.1, 0.25, 0.5\}$ in the largest and smallest patches, with local mortality in smallest patches being higher or equal to in

largest patches (Fig. S5). For each parameter combination we run 50 simulations (i.e., 5 replicate landscapes x 10 replicate communities) resulting in a total of 6,300 simulations. We completed our exploration by running the same set of simulations but with positive slopes between mortality rate and patch size (i.e., higher mortality in downstream than in upstream patches, see Appendix B and Figs. S7 & S8). This allowed us to explore and contrast biodiversity patterns with our initial predictions that are based on higher perturbation upstream (negative mortality-patch size slope). In the case of a positive slope, mortality is stronger in largest patches potentially owing to increasing human densities in downstream sites.

Experimental test

To empirically test the main prediction from our simulations, that is, patch size-dependent mortality leading to a reversal of the ‘classical’ biodiversity pattern in dendritic network, we conducted a protist metacommunity experiment. We did not experimentally test for the ‘classical’ diversity pattern without disturbances because it has already been empirically verified many times in the literature [2,3,8,26,43,44], and in the same experimental conditions [4,18]. We focussed here on testing specific predictions from the simulations owing to patch-size dependent mortality and the occurrence of the predicted reversed pattern of biodiversity under this scenario. In that context, the experiment constitutes a very conservative test of our much more general simulations. The experiment consisted of seven protist species interacting and dispersing for 29 days along 4 different dendritic networks of 36 patches (same as four of the five landscapes used in the simulations; see Fig. 1). Each landscape had 4 different patch size levels (7.5, 13, 22.5, and 45 mL) connected by dispersal along a dendritic network and preserving the

scaling properties observed in real river systems (see Fig. 1 and Fig. S9 for a photo, and section “model simulation” above, following Carrara *et al.*, 2014 [4]).

Our communities were composed of six bacterivorous protist and one rotifer species (henceforth called “protists”): *Tetrahymena* sp., *Paramecium caudatum*, *Colpidium striatum*, *Spirostomum* sp., and *Chilomonas* sp., *Blepharisma* sp. and the rotifer *Cephalodella* sp. The latter two species can, next to feeding on bacteria, to a lesser degree also predate on smaller protists. These protists were feeding on a common pool of bacteria (*Serratia fonticola*, *Bacillus subtilis* and *Brevibacillus brevis*). Prior to the beginning of the experiment, each protist species was grown in monoculture in a solution of pre-autoclaved standard protist pellet medium (Carolina Biological Supply, Burlington NC, USA, 0.46 g protist pellets 1 L⁻¹ tap water) and 10% bacteria inoculum, until they reached carrying capacity (for methodological details and protocols see Altermatt *et al.*, 2015 [38]). Protist abundance and diversity were measured by video recording combined with a trained algorithm to differentiate each species based on their morphological traits (see below).

Each microcosm in our four landscapes consisted of a 50 mL polypropylene Falcon tube (VWR, Dietikon, Switzerland). At day 0, we pipetted an equal mixture of each of the seven species into each microcosm to reach the corresponding volume (7.5, 13, 22.5 or 45 mL). Thus, protist communities were added at 15% of their carrying capacity and were allowed to grow 24 hours before the first dispersal event. Dispersal and imposed local mortality (see below) occurred two times per week, while sampling of the communities for species count was done once a week (two dispersal events between each sampling with at least 48 hours between the last dispersal/mortality event and sampling).

Sampling events and counting were done at day 0, 7, 15, 21, 29 of the experiment, while dispersal and mortality events occurred at day 1, 4, 8, 11, 16, 19, 22, 25 of the experiment.

Dispersal was done by pipetting a fixed volume (1 mL) from one patch to each of the connected patches. Dispersal was bi-directional along each edge (1 mL from a to b and 1 mL from b to a), which ensured the maintenance of the same volume in each patch throughout the duration of the experiment. We implemented bi-directional dispersal to avoid the logistical challenge of maintaining equal patch volumes with directional dispersal. To confirm that this experimental assumption did not affect our results, we conducted a sensitivity analysis with the model including an extensive number of additional simulations that demonstrate the robustness of our results to such directional dispersal (see Appendix A and Figs. S2 & S3). For dispersal we used a mirror landscape (following methods developed in Carrara *et al.*, 2012 [18]): first, 1 mL was sampled for each edge connecting a microcosm in the real landscape and then pipetted to the recipient microcosm, but in the mirror landscape. Once this was done for all microcosms, the content of each mirror microcosm was poured to the same microcosm in the real landscape. It is noteworthy, however, that because we started our simulations and experiment with species already assembled (all species were present at start in all patches), we probably are underestimating the importance of dispersal dynamics in the process of early assembly [45].

Patch size-dependent mortality was experimentally reproduced by sampling, killing and pouring back (no change in biomass) a fixed volume (1 mL) of each community regardless of patch size, which resulted in a gradient of proportionally decreasing

mortality from smaller upstream to larger downstream microcosms (this 1 mL corresponds to 13%, 8%, 4%, and 2% of disturbance-induced mortality in the respective patch sizes), and can be seen as a general type of disturbance that is blind to species identity (e.g., habitat destruction, heavy pollution). More specifically, for each mortality event, 1 mL was sampled from each microcosm and microwaved until boiling to turn all living cells into detritus [46]. After a 1-hour cooling period at ambient temperature (20 °C), the microwaved sample had reached ambient temperature again and was poured back into the same microcosm.

At each measurement day, 0.4 mL was sampled from each microcosm for the protist density measurements. Sampling was done semi-destructively, such that the sampled volume was not returned back to the patch to avoid cross-contaminations. Therefore, sampling was acting analogously to the implemented disturbance regime. Protist density was measured by using a standardized video recording and analysis procedure [47,48]. In short, a constant volume (34.4 μ L) of each 0.4 mL sample was measured under a dissecting microscope connected to a camera for the recording of videos (5 s per video, see Appendix B for further details on this method). Then, using a customized version of the R-package *bemovi* [48], we used an image processing software (ImageJ, National Institute of Health, USA) to extract the number of moving organisms per video frame along with a suite of different traits for each occurrence (e.g., speed, shape, size) that could then be used to filter out background movement noise (e.g., particles from the medium) and to identify species in a mixture (see Appendix C).

Statistical analysis

To test for the effect of patch size on protist local diversity we used a two-way linear mixed effect model testing the interactive effects of patch size and continuous time on protist species richness. To control for temporal pseudo-replication and temporal autocorrelation, we added replicate and time as nested random factors. The model was fitted by maximizing the restricted log-likelihood (“REML”, see Pinheiro *et al.*, 2016 [49]). The linear mixed effect model was conducted using the R-package NLME [49]. The complete results can be found in Table S2.

Results

Shift in diversity patterns

In the simulations without disturbance (i.e., local mortality), the highest diversity levels are found in the large downstream patches (Figs. 1 and 2a). By contrast, the smaller, mostly upstream patches have lower diversity levels. This difference in diversity unfolded with smaller patches supporting smaller population sizes (Fig. 3a), which are then more prone to extinctions relative to populations in larger patches (Fig. 3b).

In the simulations with disturbance (i.e., local mortality), highest diversity levels are found in upstream patches (Figs. 1 and 2b) consistently across all the different community (different species traits) and landscape (different spatial networks) replicates. The same pattern was found in the context of our experiment with an aquatic protist community (Figs. 1 and 2c). As in the simulations (Fig. S4), the biodiversity pattern appeared progressively over time with no apparent effects of patch size on the first sampling day (experiment day 7, 7.4 ± 0.67 species in smallest vs. 7.1 ± 0.70 species in

largest patches, mean \pm sd, Fig. S10 and Table S2), but by the final experimental day 29 there was a clear and significant decline in local diversity with patch size (4.7 ± 1.3 species in smallest patches vs. 2.5 ± 1.5 species in largest patches, mean \pm sd, Figs. 1, 2b and Table S2). Those changes in species richness were mirrored by changes in community structure (Figure S11). While being homogeneous in species number across patch sizes (see above), communities at day 7 also tended to be dominated by the fast growing species *Cephalodella* sp. (rotifer) and *Chilomonas* sp. (Figure S11). By the end of the experiment at day 29, communities were much more variable in composition but with the competitor *Paramecium caudatum* being most often dominant (Figure S11). Communities in more disturbed patches (smaller patches) were also more even in relative abundances than in larger patches where fewer species were generally dominated by a single species (Figure S11). In terms of underlying mechanisms, the simulations show that, overall, patch size-dependent mortality slows down ecological dynamics (competitive exclusion; Fig. 3a), but does more so in smaller patches (Fig. 3a), effectively increasing the time to extinction of less competitive species in those smaller patches (Fig. 3b).

Persistence of diversity patterns in river-like landscapes

Our model predicts both diversity scenarios to occur within specific parts of the parameter space defined by the magnitude of dispersal and the strength of disturbance (i.e., intercept [mortality upstream] and slope [rate of change in mortality with increasing patch size] of the relationship between patch size and local mortality rate, see Methods for details; Fig. 4). For a negative slope (higher mortality upstream), weak patch size-dependent mortality, with low intercept or slope, generates the classical diversity pattern,

while intermediate to high strengths of patch size-dependent mortality, with intermediate intercept and high slope, generate the reversed diversity pattern until local mortality is so high in small patches (high intercept) that all species upstream go extinct (see Fig. 4, $m_{upstream} = 0.5$), leading to a return to the classical diversity pattern. As expected, regardless of the strength of patch size-dependent local mortality, increasing dispersal only blurs diversity patterns by homogenizing diversity in the landscape (Figs. 4 and S6). Additional sensitivity analysis showed that the reversed biodiversity pattern is robust to strong directionality in dispersal (90 or 99% downstream, Figs. S1 and S2) because competitive exclusion still occurs later in small than in large patches. Only the combination of both high dispersal rate and strong downstream directionality can weaken the reversed biodiversity pattern (Fig. S3). Finally, a positive patch size-mortality rate relationship (higher mortality downstream than upstream) strengthens the classical diversity pattern until mortality rates are so high in larger patches that species goes extinct faster downstream relative to upstream, making the reversed pattern emerge (see Fig. S8, $m_{downstream} = 0.5$).

Discussion

Testing the impact of disturbance-induced mortality on the distribution of biodiversity in river-like landscapes, we found that disturbances can alter the classical biodiversity pattern of higher diversity in larger downstream patches predicted by the Theory of Island Biogeography ('TIB') [3,7,21]: the classical pattern persists only until a certain level of disturbance, at which point asymmetry in population turnover between smaller upstream and larger downstream patches will reverse the pattern completely. Such a reversed diversity pattern has been observed in case studies from multiple natural river systems

[9,10] and has challenged the idea of one overarching and universal diversity pattern to be expected in river-like landscapes as proposed by some studies [3,18,50,51]. However, this reversal has hitherto neither been understood mechanistically nor predicted by theory. Here, we identify a general mechanism responsible for the observed reversed diversity patterns based on theoretical considerations, which we then successfully tested and validated experimentally. The specific mechanism identified (patch-size dependent mortality) represents a general type of disturbance. However, any mechanism that induces differences in the speed of ecological dynamics between upstream and downstream patches should cause a reversal of diversity pattern. It could be differences in temperature, for instance, with lower temperatures at higher elevation slowing down upstream dynamics, or strong anthropogenic disturbance downstream (e.g., heavy pollution) which would speed up species extinction rather than slows down competitive exclusion, in large compared to small patches (see Figs S8, $m_{upstream} = 0.5$). Moreover, our approach using five different realizations of river-like networks that follow the invariant scaling properties of real riverine networks, allows us to generalize those results and their implications to riverine networks regardless of context-specific differences.

Our work indicates that the reversed pattern generated by patch-size dependent mortality is transient over time because the perturbations slow down ecological selection in smaller patches, but do not change endpoints equilibria. Similar results, related to the transient effect of disturbance, were found by [30]. Our results follow the trend by which communities are dominated early on by smaller, less competitive and fast growing species (here *Cephalodella* sp. and *Chilomonas* sp.), which are then replaced by larger and more competitive species (here *Paramecium caudatum*). The reversed diversity

pattern emerges because, across the landscape, patches are organized along a gradient in disturbance, which effectively leads to smaller upstream and larger confluence patches being at different points in their ecological selection dynamics. As selection is weaker upstream, the downstream patches reach their low diversity equilibrium more quickly. This pattern is exacerbated by highly competitive species dominating in downstream patches due to reduced disturbance, which additionally speeds up the dynamics of ecological selection locally. Consequently, it is the perturbation pattern, along with the invariant patch-size distribution in the landscape, which creates the necessary conditions for the reversed diversity gradient to emerge. We expect this effect to be especially pronounced in communities with strong competitive asymmetries and even enhanced in systems with competition-colonization trade-offs (i.e., when a weak competitor can regionally co-exist with a strong competitor because of better dispersal capacity; see [52]), as few strong competitors with low dispersal rates would lead to strong selection in the absence of turnover. Such selective processes could also affect evolutionary dynamics: Dendritic networks have recently been found to inherently lead to the emergence of non-trivial abundance patterns and eco-evolutionary dynamics per se [53,54]. Thus, the network structure will impose selection gradients, likely linked to dispersal [55,56]. In addition, patch-size dependent mortality leads to another, possibly antagonistic, selection gradient, which could lead to the emergence of non-trivial evolutionary dynamics in such networks, as has been already theoretically and empirically proposed [57]. While our study does not allow to study evolutionary dynamics (we did not consider intraspecific variation), we suggest this to be an interesting future research direction.

Disturbance-mediated slowing down of competitive exclusion is a well-understood ecological mechanism affecting population dynamics [58–61]. Here we show that this mechanism can generate a specific transient signature in terms of biodiversity patterns when applied over a spatial gradient of disturbance, which is a common feature of river-like landscapes. For instance, headwaters in river systems are naturally more prone to higher levels of disturbance than larger downstream patches because of their higher surface area exposed to disturbance relative to water volume [62,63]. Higher benthic surface area to water volume ratios are generally more net heterotrophic and thus show higher sensitivity to change in litter composition from terrestrial systems [64,65], and are also likely more sensitive to drought events because they are shallower. They are on average also located at higher elevation, and thus experiencing stronger variations in temperature and weather conditions. Human-induced disturbances in pristine headwaters are increasingly common [58] and the general mechanisms demonstrated by our study suggest that these anthropogenic impacts can have unexpected consequences by inverting large-scale biodiversity patterns. Although more robust to change, our modeling results also suggest that very high disturbance levels in downstream locations can also lead to a reversed diversity pattern. This is likely relevant around large urban centers that are generally located at downstream confluences.

In conclusion, we show that patch size-dependent mortality (whether negative or positive), which is commonly displayed in river-like landscapes, can be a key mechanism responsible for shaping the distribution of biodiversity distribution in riverine landscapes.

428 **Acknowledgement**

429 We thank S. Gut, S. Flückiger and E. Keller for help during the laboratory work.

430 Funding is from the Swiss National Science Foundation Grants PP00P3_150698 and

431 PP00P3_179089 to FA.

References

1. Vannote RL, Minshall GW, Cummins KW, Sedell JR, Cushing CE. 1980 The river continuum concept. *Can. J. Fish. Aquat. Sci.* **37**, 130–137. (doi:10.1139/f80-017)
2. Ward JV, Tockner K, Arscott DB, Claret C. 2002 Riverine landscape diversity. *Freshw. Biol.* **47**, 517–539. (doi:10.1046/j.1365-2427.2002.00893.x)
3. Muneeppeerakul R, Bertuzzo E, Lynch HJ, Fagan WF, Rinaldo A, Rodriguez-Iturbe I. 2008 Neutral metacommunity models predict fish diversity patterns in Mississippi–Missouri basin. *Nature* **453**, 220–222. (doi:10.1038/nature06813)
4. Carrara F, Rinaldo A, Giometto A, Altermatt F, Roelke AEDL, Kalisz ES. 2014 Complex interaction of dendritic connectivity and hierarchical patch size on biodiversity in river-like landscapes. *Am. Nat.* **183**, 13–25. (doi:10.1086/674009)
5. Deiner K, Fronhofer EA, Mächler E, Walser J-C, Altermatt F. 2016 Environmental DNA reveals that rivers are conveyor belts of biodiversity information. *Nat. Commun.* **7**, 12544. (doi:10.1038/ncomms12544)
6. Muneeppeerakul R, Weitz JS, Levin SA, Rinaldo A, Rodriguez-Iturbe I. 2007 A neutral metapopulation model of biodiversity in river networks. *J. Theor. Biol.* **245**, 351–363. (doi:10.1016/j.jtbi.2006.10.005)
7. Economo EP, Keitt TH. 2010 Network isolation and local diversity in neutral metacommunities. *Oikos* **119**, 1355–1363. (doi:10.1111/j.1600-0706.2010.18272.x)
8. Seymour M, Fronhofer EA, Altermatt F. 2015 Dendritic network structure and dispersal affect temporal dynamics of diversity and species persistence. *Oikos* **124**, 908–916. (doi:10.1111/oik.02354)
9. Besemer K, Singer G, Quince C, Bertuzzo E, Sloan W, Battin TJ. 2013 Headwaters are critical reservoirs of microbial diversity for fluvial networks. *Proc R Soc B* **280**, 20131760. (doi:10.1098/rspb.2013.1760)
10. Tornwall BM, Swan CM, Brown BL. 2017 Manipulation of local environment produces different diversity outcomes depending on location within a river network. *Oecologia* **184**, 663–674. (doi:10.1007/s00442-017-3891-7)
11. Balian EV, Segers H, Lévêque C, Martens K. 2008 The Freshwater Animal Diversity Assessment: an overview of the results. *Hydrobiologia* **595**, 627–637. (doi:10.1007/s10750-007-9246-3)
12. Tockner K, Stanford JA. 2002 Riverine flood plains: present state and future trends. *Environ. Conserv.* **29**, 308–330. (doi:10.1017/S037689290200022X)
13. Dudgeon D *et al.* 2006 Freshwater biodiversity: importance, threats, status and conservation challenges. *Biol. Rev.* **81**, 163–182. (doi:10.1017/S1464793105006950)

- 467 14. Vörösmarty CJ *et al.* 2010 Global threats to human water security and river
468 biodiversity. *Nature* **467**, 555–561. (doi:10.1038/nature09440)
- 469 15. Rodriguez-Iturbe I, Muneeppeerakul R, Bertuzzo E, Levin SA, Rinaldo A. 2009 River
470 networks as ecological corridors: A complex systems perspective for integrating
471 hydrologic, geomorphologic, and ecologic dynamics. *Water Resour. Res.* **45**,
472 W01413. (doi:10.1029/2008WR007124)
- 473 16. Rodriguez-Iturbe I, Rinaldo A. 1997 *Fractal river basins: chance and self-*
474 *organization*. New York, N.Y.: Cambridge University Press.
- 475 17. Mari L, Casagrandi R, Bertuzzo E, Rinaldo A, Gatto M. 2014 Metapopulation
476 persistence and species spread in river networks. *Ecol. Lett.* **17**, 426–434.
477 (doi:10.1111/ele.12242)
- 478 18. Carrara F, Altermatt F, Rodriguez-Iturbe I, Rinaldo A. 2012 Dendritic connectivity
479 controls biodiversity patterns in experimental metacommunities. *Proc. Natl. Acad.*
480 *Sci.* **109**, 5761–5766. (doi:10.1073/pnas.1119651109)
- 481 19. Leopold LB, Wolman MG, Miller JP. 1995 *Fluvial Processes in Geomorphology*.
482 Trade Paperback Edition edition. New York: Dover Publications.
- 483 20. Leibold MA *et al.* 2004 The metacommunity concept: a framework for multi-scale
484 community ecology. *Ecol. Lett.* **7**, 601–613. (doi:10.1111/j.1461-0248.2004.00608.x)
- 485 21. MacArthur RH, Wilson EO. 1963 An equilibrium theory of insular zoogeography.
486 *Evolution* **17**, 373. (doi:10.2307/2407089)
- 487 22. MacArthur RH, Wilson EO. 1967 *The Theory of Island Biogeography*. Reprint
488 edition. Princeton: Princeton University Press.
- 489 23. Hubbell SP. 2001 *The Unified Neutral Theory of Biodiversity and Biogeography*.
490 Princeton University Press.
- 491 24. Angela DY, Lei SA. 1998 Equilibrium Theory of Island Biogeography: A Review.
492 *Proc. RMRS* , 163.
- 493 25. Harvey E, MacDougall AS. 2014 Trophic island biogeography drives spatial
494 divergence of community establishment. *Ecology* **95**, 2870–2878. (doi:10.1890/13-
495 1683.1)
- 496 26. Finn DS, Bonada N, Múrria C, Hughes JM. 2011 Small but mighty: headwaters are
497 vital to stream network biodiversity at two levels of organization. *J. North Am.*
498 *Benthol. Soc.* **30**, 963–980. (doi:10.1899/11-012.1)
- 499 27. Lorenz S, Rasmussen JJ, Süß A, Kalettka T, Golla B, Horney P, Stähler M, Hommel
500 B, Schäfer RB. 2017 Specifics and challenges of assessing exposure and effects of

pesticides in small water bodies. *Hydrobiologia* **793**, 213–224. (doi:10.1007/s10750-016-2973-6)

28. Palmer MA *et al.* 2010 Mountaintop mining consequences. *Science* **327**, 148–149. (doi:10.1126/science.1180543)

29. Valladares F, Bastias CC, Godoy O, Granda E, Escudero A. 2015 Species coexistence in a changing world. *Front. Plant Sci.* **6**. (doi:10.3389/fpls.2015.00866)

30. Violle C, Pu Z, Jiang L. 2010 Experimental demonstration of the importance of competition under disturbance. *Proc. Natl. Acad. Sci.* **107**, 12925–12929. (doi:10.1073/pnas.1000699107)

31. McCann KS. 2011 *Food Webs*. Princeton University Press.

32. Hemphill N. 1991 Disturbance and variation in competition between two stream insects. *Ecology* **72**, 864–872. (doi:10.2307/1940588)

33. Townsend CR, Scarsbrook MR, Dolédec S. 1997 The intermediate disturbance hypothesis, refugia, and biodiversity in streams. *Limnol. Oceanogr.* **42**, 938–949. (doi:10.4319/lo.1997.42.5.0938)

34. Shea K, Roxburgh SH, Rauschert ESJ. 2004 Moving from pattern to process: coexistence mechanisms under intermediate disturbance regimes. *Ecol. Lett.* **7**, 491–508. (doi:10.1111/j.1461-0248.2004.00600.x)

35. Urban MC. 2004 Disturbance Heterogeneity Determines freshwater metacommunity structure. *Ecology* **85**, 2971–2978. (doi:10.1890/03-0631)

36. Ledger ME, Harris RML, Armitage PD, Milner AM. 2008 Disturbance frequency influences patch dynamics in stream benthic algal communities. *Oecologia* **155**, 809–819. (doi:10.1007/s00442-007-0950-5)

37. Srivastava DS *et al.* 2004 Are natural microcosms useful model systems for ecology? *Trends Ecol. Evol.* **19**, 379–84. (doi:10.1016/j.tree.2004.04.010)

38. Altermatt F *et al.* 2015 Big answers from small worlds: a user’s guide for protist microcosms as a model system in ecology and evolution. *Methods Ecol. Evol.* **6**, 218–231. (doi:10.1111/2041-210X.12312)

39. Thorp JH, Thoms MC, DeLong MD. 2006 The riverine ecosystem synthesis: biocomplexity in river networks across space and time. *River Res. Appl.* **22**, 123–147. (doi:10.1002/rra.901)

40. Giometto A, Rinaldo A, Carrara F, Altermatt F. 2013 Emerging predictable features of replicated biological invasion fronts. *Proc. Natl. Acad. Sci.* **201321167**. (doi:10.1073/pnas.1321167110)

- 535 41. Rigon R, Rinaldo A, Rodriguez-Iturbe I, Bras RL, Ijjasz-Vasquez E. 1993 Optimal
536 channel networks: A framework for the study of river basin morphology. *Water*
537 *Resour. Res.* **29**, 1635–1646. (doi:10.1029/92WR02985)
- 538 42. Rinaldo A, Banavar JR, Maritan A. 2006 Trees, networks, and hydrology. *Water*
539 *Resour. Res.* **42**, W06D07. (doi:10.1029/2005WR004108)
- 540 43. Altermatt F, Seymour M, Martinez N. 2013 River network properties shape α -
541 diversity and community similarity patterns of aquatic insect communities across
542 major drainage basins. *J. Biogeogr.* **40**, 2249–2260. (doi:10.1111/jbi.12178)
- 543 44. Altermatt F, Alther R, Fišer C, Jokela J, Konec M, Kury D, Mächler E, Stucki P,
544 Westram AM. 2014 Diversity and distribution of freshwater amphipod species in
545 Switzerland (Crustacea: Amphipoda). *PLoS ONE* **9**, e110328.
546 (doi:10.1371/journal.pone.0110328)
- 547 45. Walker LR, Chapin FS. 1987 Interactions among processes controlling successional
548 change. *Oikos* **50**, 131–135. (doi:10.2307/3565409)
- 549 46. Harvey E, Gounand I, Ganesanandamoorthy P, Altermatt F. 2016 Spatially cascading
550 effect of perturbations in experimental meta-ecosystems. *Proc R Soc B* **283**,
551 20161496. (doi:10.1098/rspb.2016.1496)
- 552 47. Pennekamp F, Schtickzelle N. 2013 Implementing image analysis in laboratory-based
553 experimental systems for ecology and evolution: a hands-on guide. *Methods Ecol.*
554 *Evol.* **4**, 483–492. (doi:10.1111/2041-210X.12036)
- 555 48. Pennekamp F, Schtickzelle N, Petchey OL. 2015 BEMOVI, software for extracting
556 behavior and morphology from videos, illustrated with analyses of microbes. *Ecol.*
557 *Evol.* **5**, 2584–2595. (doi:10.1002/ece3.1529)
- 558 49. Pinheiro J, Bates D, DebRoy S, Sarkar D, R Core Team. 2016 *nlme: Linear and*
559 *Nonlinear Mixed Effects Model*. See <http://CRAN.R-project.org/package=nlme>.
- 560 50. Widder S, Besemer K, Singer GA, Ceola S, Bertuzzo E, Quince C, Sloan WT,
561 Rinaldo A, Battin TJ. 2014 Fluvial network organization imprints on microbial co-
562 occurrence networks. *Proc. Natl. Acad. Sci.* **111**, 12799–12804.
563 (doi:10.1073/pnas.1411723111)
- 564 51. Bertuzzo E, Maritan A, Gatto M, Rodriguez-Iturbe I, Rinaldo A. 2007 River
565 networks and ecological corridors: Reactive transport on fractals, migration fronts,
566 hydrochory. *Water Resour. Res.* **43**, W04419. (doi:10.1029/2006WR005533)
- 567 52. Cadotte MW. 2007 Competition–colonization trade-offs and disturbance effects at
568 multiple scales. *Ecology* **88**, 823–829. (doi:10.1890/06-1117)

- 569 53. Fronhofer EA, Altermatt F. 2017 Classical metapopulation dynamics and eco-
570 evolutionary feedbacks in dendritic networks. *Ecography* **40**, 1455–1466.
571 (doi:10.1111/ecog.02761)
- 572 54. Altermatt F, Fronhofer EA. 2018 Dispersal in dendritic networks: Ecological
573 consequences on the spatial distribution of population densities. *Freshw. Biol.* **63**,
574 22–32. (doi:10.1111/fwb.12951)
- 575 55. Tonkin JD, Altermatt F, Finn DS, Heino J, Olden JD, Pauls SU, Lytle DA. 2018 The
576 role of dispersal in river network metacommunities: Patterns, processes, and
577 pathways. *Freshw. Biol.* **63**, 141–163. (doi:10.1111/fwb.13037)
- 578 56. Anderson KE, Hayes SM. 2018 The effects of dispersal and river spatial structure on
579 asynchrony in consumer–resource metacommunities. *Freshw. Biol.* **63**, 100–113.
580 (doi:10.1111/fwb.12998)
- 581 57. Paz-Vinas I, Loot G, Stevens VM, Blanchet S. 2015 Evolutionary processes driving
582 spatial patterns of intraspecific genetic diversity in river ecosystems. *Mol. Ecol.* **24**,
583 4586–4604. (doi:10.1111/mec.13345)
- 584 58. Sheil D. 2016 Disturbance and distributions: avoiding exclusion in a warming world.
585 *Ecol. Soc.* **21**. (doi:10.5751/ES-07920-210110)
- 586 59. Chesson PL, Warner RR. 1981 Environmental variability promotes coexistence in
587 lottery competitive systems. *Am. Nat.* **117**, 923–943.
- 588 60. Chesson P. 2000 Mechanisms of maintenance of species diversity. *Annu. Rev. Ecol.*
589 *Syst.* **31**, 343–366.
- 590 61. Göthlich L, Oschlies A. 2015 Disturbance characteristics determine the timescale of
591 competitive exclusion in a phytoplankton model. *Ecol. Model.* **296**, 126–135.
592 (doi:10.1016/j.ecolmodel.2014.10.033)
- 593 62. Lowe WH, Likens GE, Power ME. 2006 Linking Scales in Stream Ecology.
594 *BioScience* **56**, 591. (doi:10.1641/0006-3568(2006)56[591:LSISE]2.0.CO;2)
- 595 63. Battin TJ, Kaplan LA, Findlay S, Hopkinson CS, Marti E, Packman AI, Newbold JD,
596 Sabater F. 2008 Biophysical controls on organic carbon fluxes in fluvial networks.
597 *Nat. Geosci.* **1**, 95–100. (doi:10.1038/ngeo101)
- 598 64. England LE, Rosemond AD. 2004 Small reductions in forest cover weaken
599 terrestrial-aquatic linkages in headwater streams. *Freshw. Biol.* **49**, 721–734.
600 (doi:10.1111/j.1365-2427.2004.01219.x)
- 601 65. Rasmussen JB. 2010 Estimating terrestrial contribution to stream invertebrates and
602 periphyton using a gradient-based mixing model for $\delta^{13}\text{C}$. *J. Anim. Ecol.* **79**, 393–
603 402. (doi:10.1111/j.1365-2656.2009.01648.x)

Figure legends

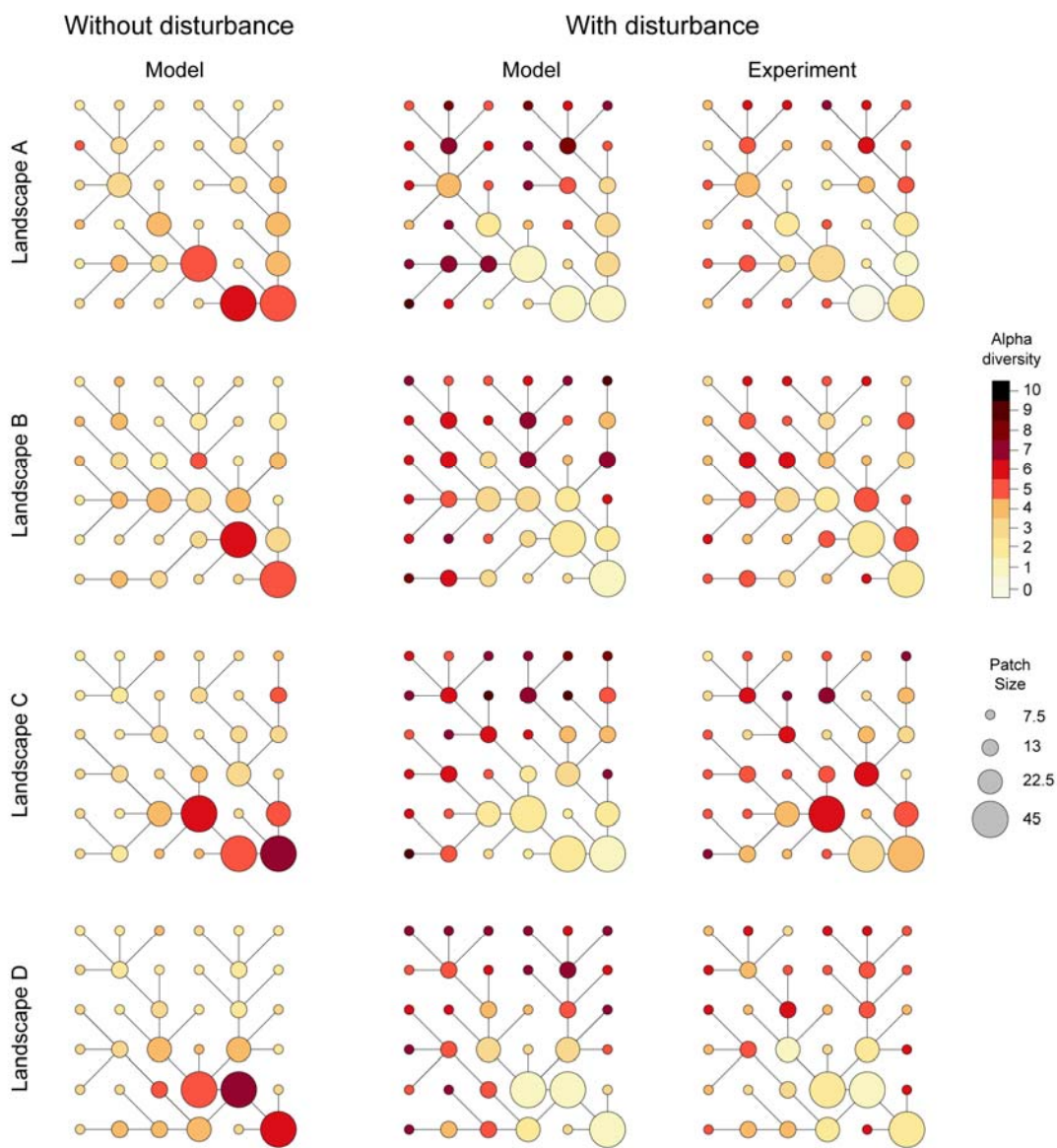
Figure 1. Diversity distribution (local species richness) in dendritic, river-like landscapes with and without disturbance. The ‘Model’ landscapes displayed in this figure are examples of simulations made with the same replicate community. Results were qualitatively similar for each of the 10 replicate communities randomly generated (see Fig. S4); model parameters specific to these simulations are dispersal $\delta_V = 0.05$, mortality rates m are 0.1, 0.0868, 0.064 and 0.01 respectively from smaller to larger patches for simulation with disturbance (patch size-dependent mortality), and 0 without. For network graphs based on experimental results, each graph represents local diversity pattern for one of the four landscape replicates (A, B, C, and D). For illustration purpose, all network graphs are depicted at snapshots of diversity patterns at time of comparable local diversity: time 118.4 (simulations without disturbance; faster dynamics), time 167.9 (simulations with disturbance; slower dynamics), and at experimental day 29 (last sampling day). See complete dynamics over time for model simulations on Fig. S4 and for experimental results on Fig. S5.

Figure 2. Diversity distribution in dendritic, river-like landscapes a) without and b) with disturbance. For the panels based on simulations, each boxplot represents the distribution of diversity values across the 4 replicate landscapes used in the experiment, and for one of the 10 replicate communities that were used in the simulations (see Fig. S4 for all 10 replicates). Parameters and data are the same as in Figure 1. The two largest patch sizes of 45 (N=11) and 22.5 mL (N=14) show significant decline in local diversity compared to the smaller patch sizes of 13 (N=34) and 7.5 mL (N=85) (see Table S1 for complete linear mixed effect model results).

Figure 3. Time to extinction as a function of disturbance in the simulations. a) Example of temporal dynamics of one species in each patch of one landscape with and without disturbance (dotted and full lines respectively). Arrows indicate time at extinction for the smallest (red) and largest (blue) patches with and without disturbance. b) Time at extinction for each species in the smallest and the largest patches with disturbance (lower panel) or without disturbance (upper panel). Each boxplot represents the distribution of values across the patches of 4 replicate landscapes, and for one of the 10 different replicate communities that were used in the simulations (see Fig. S4 for all replicates). Data and parameters are the same as in Fig. 1.

Figure 4. Persistence of diversity patterns in dendritic, river-like landscapes. The figure illustrates the parameter space within which each diversity pattern is found as a function of the strength of disturbance (mortality rate-patch size relationship) and the magnitude of dispersal (for more dispersal rates see Fig. S6). δ_V is the dispersal volume, which we vary the dispersal rate ($d_x = \delta_V/V_x$, with V_x the volume of patch x). $\alpha_D - \alpha_U$ corresponds to the maximum observed difference, along the temporal dynamics, in richness between largest and smallest patches. When $\alpha_D - \alpha_U$ is positive (green colours) the classical pattern emerges over the course of the dynamics, with higher diversity in large than in small patches, while when the difference is negative (pink colours) it is the reversed pattern, with higher diversity in small (i.e., headwater) rather than in large patches. Each case represents an average over 50 replicate simulations (5 different landscapes x 10 different communities). Mortality rate increase in small upstream patches raises the intercept of the negative mortality rate-patch size relationship, while mortality rate increase in large downstream patches, reduces its slope.

651 **Figure 1**



652

653

654

655 **Figure 2**

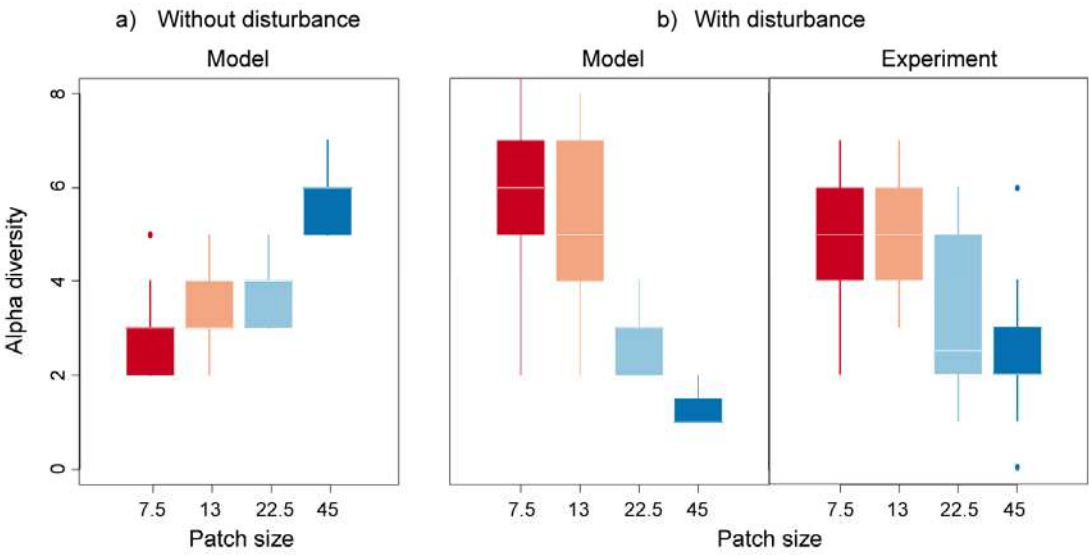
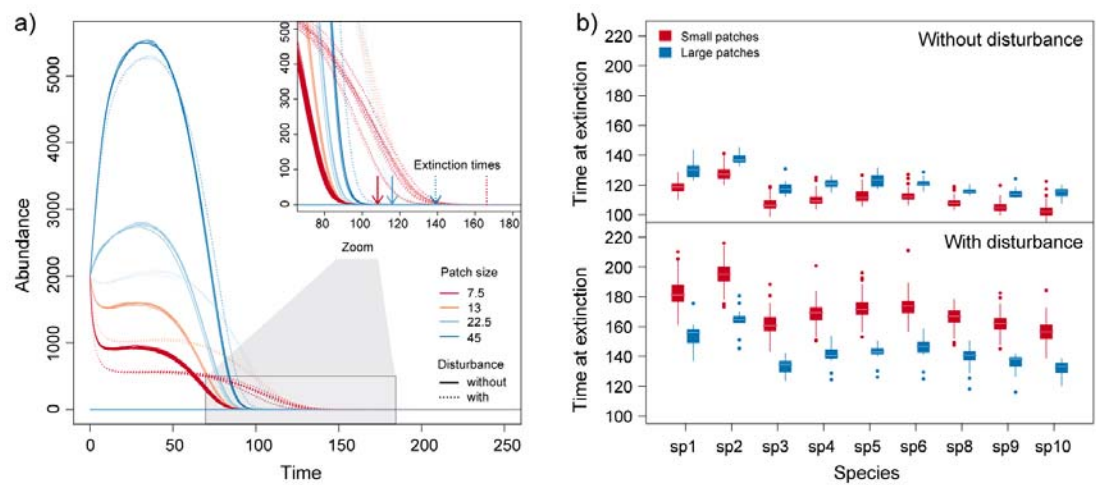
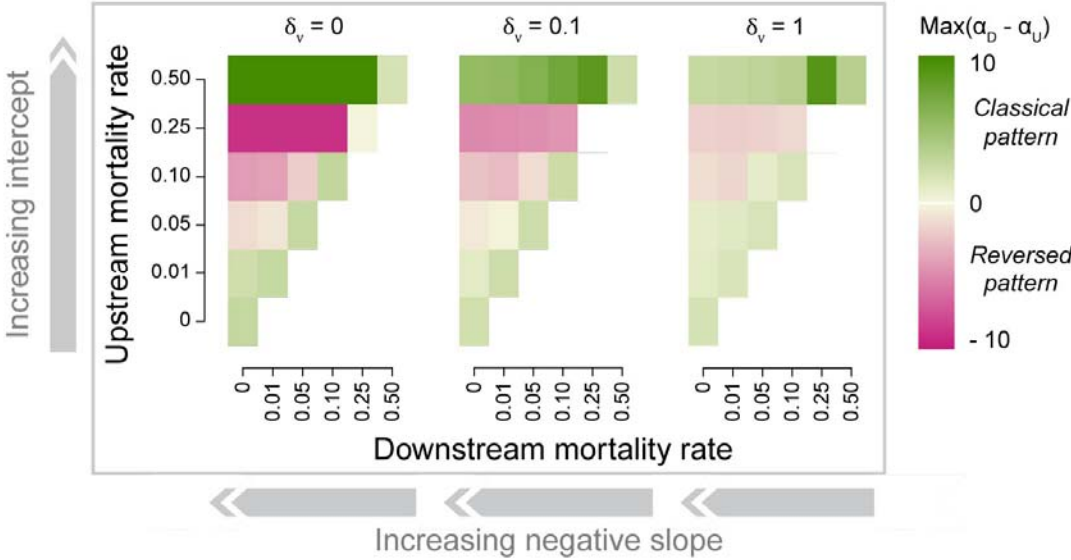


Figure 3



661 **Figure 4**



662

663